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### UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

WASHINGTON, D.C. 20460

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## OPP OPPICIAL RECORD HEALTH EFFECTS DIVISION SCIENTIFIC DATA REVIEWS EFA SERIES 361

**MEMORANDUM** 

September 24, 1998

OFFICE OF PREVENTION, PESTICIDES AND TOXIC SUBSTANCES

SUBJECT:

REVISED Oxyflourfen (Goal) Quantitative Risk Assessment (Q<sub>1</sub>) Based On CD-1 Male Mouse Dietary Study With <sup>3</sup>/<sub>4</sub>'s

Interspecies Scaling Factor

P.C. Code 111601

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Summary

The unit risk,  $Q_1^*$  (mg/kg/day)<sup>-1</sup>, of Oxyflourfen (Goal) based upon male mouse combined liver tumor (adenomas and/or carcinomas) rates is  $7.32 \times 10^{-2}$  in human equivalents (converted from animals to humans by use of the  $^3/_4$ 's scaling factor - Tox\_Risk program, Version 3.5, K. Crump, 1994)<sup>1</sup>. The dose levels used from the 20-month dietary study were 0, 2, 20, and 200 ppm of Oxyflourfen (Goal). The corresponding tumor rates for the male mouse combined liver tumors (adenomas and/or carcinomas) were 2/47, 0/44, 4/44, and 8/52, respectively.

#### Background

On May 24, 1989, the Carcinogenicity Peer Review Committee recommended that a quantitative risk assessment for Oxyflourfen (Goal) be estimated for combined liver tumors (adenomas and/or carcinomas) in male mice. A quantitative risk assessment (Oxyflourfen (Goal) - Quantitative Risk Assessment, 20 Month Dietary Study of Charles River CD Male Mice, B. Fisher, 9/18/89) was prepared using the  $^2/_3$ 's scaling factor. This revised quantitative risk assessment reflects the Division change from use of the  $^2/_3$ 's scaling factor to the  $^3/_4$ 's scaling factor in  $1994^1$ .

<sup>&</sup>lt;sup>1</sup>See memo - Deriving Q<sub>1</sub>'s Using the Unified Interspecies Scaling Factor, P.A. Fenner-Crisp, Director, HED, 7/1/94.

The statistical evaluation (Oxyflourfen (Goal) - Qualitative Risk Assessment - 20 Month Feeding Study - Charles River CD Male Mice, B. Fisher, 10/28/88) indicated that there were significant decreasing trends with either the untreated or vehicular (ethanol) control groups, but no significant pair-wise comparisons, for mortality with increasing doses of Oxyflourfen (Goal). The male mice had a dose-related significant increasing trend with either control group at p < 0.01, and a significant difference in the pair-wise comparison of the 200 ppm dose group with the vehicular controls at p < 0.05, for combined liver tumors (adenomas and/or carcinomas).

### Dose-Response Analysis

The estimate of unit risk,  $Q_1$ , was based upon combined liver tumors (adenomas and/or carcinomas) observed in male; mice.

Since the male mice had no statistically significant incremental changes in mortality with increasing doses of Oxyflourfen (Goal), the estimate of the unit risk, Q<sub>1</sub>, was obtained by the application of the Multi-Stage model (Tox\_Risk program, Version 3.5, K. Crump, 1994).

For the conversion to human equivalents, weights of 0.03 kg for the mouse, 70 kg for humans and the  $^3/_4$ 's scaling factor were used.

It is to be noted that the  $Q_1^*$  (mg/kg/day)<sup>-1</sup> is an estimate of the <u>upper bound</u> on risk and that, as stated in the EPA Risk Assessment Guidelines, "the true value of the risk is unknown, and may be as low as zero."